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(54) Title: HEPATOBLASTS AND METHOD OF ISOLATING SAME

(57) Abstract

This invention relates to methods of isolating hepatoblasts utilizing panning techniques and fluorescence activated cell sorting. This invention further relates to isolated hepatoblasts and to a method of treating liver dysfunction as well as to methods of forming artificial

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HEPATOBLASTS AND METHOD OF ISOLATING SAME CROSS-REFERENCE TO RELATED APPLICATIONS

This Application is a Continuation-In-Part of Application Serial No. 07/741,128 filed August 7, 1991, entitled PROLIFERATION OF HEPATOCYTE PRECURSORS.

FIELD OF THE INVENTION

This invention relates to methods for isolating hepatoblasts and to said isolated hepatoblasts. isolated hepatoblasts of the invention comprise liver cells (pluripotent precursors) committed and 10 progenitors (precursors with only one fate) for either cells. The isolated duct or bile hepatocytes hepatoblasts of the invention may be used to treat liver dysfunction and for artificial livers, gene therapy, 15 drug testing and vaccine production. In addition, the isolated hepatoblasts of the invention may be used for research, therapeutic and commercial purposes which require the use of populations of functional liver cells.

Unlike mature liver cells, the hepatoblasts of
the invention generate daughter cells that can mature
through the liver lineage and offer the entire range of
liver functions, many of which are lineage-position
specific. Further, the hepatoblasts of the invention
have a greater capacity for proliferation and long-term
viability than do mature liver cells. As a result, the
hepatoblasts of the invention are better for research,
therapeutic and commercial uses than mature liver cells.

BACKGROUND OF THE INVENTION

Stem cells and early progenitors have long been 30 known to exist in rapidly proliferating adult tissues such as bone marrow, gut and epidermis, but have only recently been thought to exist in quiescent tissues such as adult liver, an organ characterized by a long cellular life span. The ability of stem cells to

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self-replicate and produce daughter cells with multiple fates distinguishes them from committed progenitors. contrast, committed progenitors produce daughter cells with only one fate in terms of cell type, and these cells undergo a gradual maturation process wherein differentiated functions · appear in lineage-position-dependent process.

adult organisms, stem cells in somatic tissues produce a lineage of daughter cells that undergo a unidirectional, terminal differentiation process. well-characterized lineage systems, hemopoiesis, gut and epidermis, stem cells have been identified by empirical assays in which the stem cells were shown to be capable of producing the full range of To date, no molecular markers are known descendants. which uniquely identify stem cells as a general class of cells, and no molecular mechanisms are known which result in the conversion of cells from self-replication and pluripotency to a commitment to differentiation and a single fate.

The structural and functional units of hepatic parenchyma is the acinus, which is organized like a wheel around two distinct vascular beds. sets of portal triads, each with a portal venule, a hepatic arteriole and a bile duct, form the periphery, and the central vein forms the hub. The parenchyma, which comprises the "spokes" of the wheel, consists of plates of cells lined on both sides by the fenestrated sinusoidal endothelium. Blood flows from the portal 30 venules and hépatic arterioles at the portal triads, through sinusoids which align plates of parenchyma, to terminal hepatic venules, the central Hepatocytes display marked morphologic, biochemical and functional heterogeneity based on their acinar location (see Gebhardt, Pharmac. Ther., Vol. 53, pp. 275-354 (1990)).

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Comparatively, periportal parenchymal cells are small in size, midacinar cells are intermediate in size and pericentral cells are largest in size. acinar-position-dependent variations in the morphology 5 of mitochondria, endoplasmic reticulum and glycogen Of critical importance is that the diploid granules. and those with greatest parenchymal cells periportally. In parallel, are located potential expression tissue-specific gene acinar-position-dependent leading to the hypothesis that 10 the expression of genes is maturation-dependent (see Sigal et al., Amer. J. Physiol., Vol. 263, pp. G139-G148 (1993)).

It is currently believed that the liver is a stem cell and lineage system which has several parallels 15 to the gut, skin and hemopoietic systems (see Sigal et al., Amer. J. Physiol., Vol. 263, pp. G139-G148 (1993); Sigal et al. In Extracellular Matrix, Zern and Reed, eds, Marcel Dekker, NY., pp. 507-537 (1993); and Brill et al., Liver Biology and Pathobiology, Arias et al., 3d 20 eds, Raven Press, NY (1994 in press)). As such, it is expected that there are progenitor cell populations in the livers of all or most ages of animals. A lineage model of the liver would clarify why researches have been unable to grow adult, mature liver cells in culture 25 for more than a few rounds of division, have observed only a few divisions of mature, adult liver cells when injected in vivo into liver or into ectopic sites, and have had limited success in establishing artificial livers with adult liver cells. These impasses are of considerable concern in the use of isolated liver cells liver transplantation, artificial for therapy and other therapeutic and commercial uses.

The success of the above-listed procedures sometimes the use of hepatic progenitor cells (hepatoblasts) which are found in a high proportion of liver cells in early embryonic livers and in small

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numbers located periportally in adult livers. it is desirable to isolate such hepatoblasts, a need has arisen to develop a method of successfully isolating The inventors have identified hepatoblasts. for isolating method developed а and markers hepatoblasts from the livers of animals at any age. methods of the invention have been developed using embryonic and neonatal livers from rats, however, the method of the invention offers a systematic approach to isolating hepatoblasts from any age from any species.

invention have been the methods οf The developed with embryonic livers in which there are significant numbers of pluripotent liver cells (liver stem cells) and committed progenitors (cells with a single fate to become either hepatocytes or bile duct The onset of differentiation of rat parenchymal liver occurs by the tenth day of cells of the gestation. By this stage, parenchymal cells (epithelial or epitheloid cells) are morphologically homogeneous and consist of small cells with scant cytoplasm and, therefore, high nuclear to cytoplasmic ratios, with undifferentiated, pale, nuclei and a few intercellular adhesions. Most liver parenchymal cells at this stage are considered to be bipotent for bile duct cells and hepatocytes. Although they express, usually weakly, some liver-specific functions known to be activated very albumin such as development, early in they đο not express (AFP), α-fetoprotein adult-specific markers such as glycogen, urea-cycle enzymes or major urinary protein (MUP). Only a few islands of fetal cells are positive for BDS, a bile duct cell-specific marker, and none are positive for HES6, a hepatocyte-specific marker (see Germain et al., Cancer Research, Vol. 48, pp. 4909-4918 (1988)). cytoplasm hepatoblasts with scant The ovoid-shaped nuclei comprise several cell populations including pluripotent liver stem cells and committed

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progenitors, each having only one fate for either bile duct cells or hepatocytes.

By the fifteenth day of gestation, hepatoblasts increasingly are comprised of the committed progenitors that differentiate along either the bile duct or the hepatocytic lineage. Their maturation is denoted by in morphology (increasing size, changes organelles and vacuoles, cytoplasmic numbers ο£ heterogeneous nuclear morphologies and an increase in pigmented granules), which can be distinguished readily flow cytometric parameters. "Forward "Side scatter" measures cellular measures cell size. complexity or granularity, which is affected by the numbers of cellular organelles. Autofluorescence is dependent upon lipofuscins and other pigments that increase with maturation.

Accompanying the morphological changes step-wise or sequential changes in expression of types antigens surface cytokeratins, various tissue-specific genes. Whereas the early hepatoblasts which include liver stem cells intensely express AFP and weakly express albumin, committed progenitors destined to become hepatocytes form cords of cells that lose their AFP expression, express increasingly high levels albumin and gradually acquire hepatocyte-specific markers such as glycogen and urea cycle enzymes. Cells destined to become intrahepatic bile duct cells arise hepatoblasts and seemingly identical expression of AFP, lose albumin expression and acquire string of Initially, а (CK 19). 19 cytokeratin pearl-like cells is present around the large vascular branches close to the liver hilium. Over the ensuing days, similar structures appear throughout the liver. BDS7-positive cells rapidly enlarge and become more numerous with increasing developmental age. Gradually, lumina form within the structures, and by the eighteenth day of gestation, bile ductular structures are morphologically identifiable.

In order to understand liver development the sequential changes in the expression liver-specific genes with maturation, it is necessary to study the hepatoblasts directly. However, the study of hepatoblasts is hindered by the difficulty in isolating them since they always constitute a small portion, less 10%, of the cell types within the liver embryonic, neonatal, and adult life. 10 In the embryo, the liver is the site for both hepatopoiesis (formation of cells) and hemopoiesis (formation cells). Hempoietic cells migrate from the yolk sac into liver the during the twelfth day of gestation. Subsequently, hemopoiesis, particularly erythropoiesis, 15 rapidly becomes one of the most prominent functions of the fetal liver with hemopoietic cells comprising 50% or more of the liver mass. In neonates, the majority of the liver cells are either hemopoietic cells or mature 20 liver cells (hepatocytes or bile duct cells). result, sequential changes in parenchymal functions in intact liver are difficult to interpret because the data are confounded by the changing hemopoietic For example, it has been demonstrated contributions. that a transient decrease in parenchymal functions at 25 day eighteen of gestation is due not to a decrease in hepatic cells or in their expression of these genes, but occurs because it is the peak of erythropoiesis, when the liver consists of erythroid Hemopoiesis in the liver declines rapidly after birth as it transfers to the bone marrow, the site of hemopoiesis in the adult. Nevertheless, isolation of hepatoblasts in adult liver remains problematic, since they comprise a very small percentage of hepatic cells.

35 Because hepatoblasts can generate all developmental stages of liver cells and, therefore, offer the entire range of liver-specific functions

encoded by genes activated and expressed in early to late stages of differentiation, have much greater growth potential than mature liver cells, have greater proliferative potential and offer cells with greater ability for transfection with appropriate genes (i.e., greater capacity for gene therapy), it is desirable to isolate hepatoblasts (as opposed to mature liver cells).

Currently available methods for isolation of hepatoblasts require the use of fractionation methods 10 for cell size or cell density which are inadequate for separating the hemopoietic from hepatopoietic the precursors, require the use of cells surviving specific enzyme treatments such as pronase digestion (which have been proven to also kill hepatoblast subpopulations) or 15 require the use of selection protocols in culture in which enrichment of the cells of interest are dependent upon differential attachment to the substratum or differential growth in specific culture media. Hence, currently available isolation methods have proven very Moreover, identification 20 inefficient. parenchymal cell precursors is dependent upon assays for parenchymal-specific functions. Further, hepatoblasts most culture conditions dedifferentiate under thereby come undetectable, or there are such a high 25 proportion of non-relevant cells (e.g., mesenchymal cells) that the functions of interest are swamped out by those of the contaminant cell populations. In addition, dissociated liver cells readily from large aggregates calciumand temperature-dependent 30 glycoprotein-mediated process. In order to disaggregate the liver cells, it is necessary to utilize mechanical methods including vigorous pipetting and aspiration methods which are syringe, through insufficient to achieve single cell suspensions and 35 which can result in dramatically reduced viability of the cells. Hence it is desirable to develop a method of isolating fetal hepatoblasts which method maintains the

hepatoblasts as a single cell suspension, does not result in cell aggregation, and is applicable to all ages.

It is therefore an object of this invention to provide methods of isolating hepatoblasts.

It is a further object of this invention to provide isolated hepatoblasts.

It is another object of this invention to provide a method of utilizing isolated hepatoblasts to treat liver dysfunction.

It is a still further object of this invention to provide methods of forming artificial livers utilizing isolated hepatoblasts.

SUMMARY OF THE INVENTION

This invention relates to isolated hepatoblasts 15 and to methods of isolating hepatoblasts utilizing flow cytometry (fluorescence panning techniques and activated cell sorting) on cell suspensions of liver cells Dissociated liver are panned cells. 20 fluorescence activated cell sorted utilizing antibodies so as to greatly reduce the numbers of contaminating cell types, such as hemopoietic cells in embryonic liver or mature liver cells in adults. The cells that do not adhere to the panning dishes are negatively sorted using multiple antibodies to the contaminant cell types which 25 leads to a cell population highly enriched for immature hepatic cell types, and then segregated into distinct immature hepatic cell types subcategories of fluorescence activated cell multiparametric sorting. 30 This invention is further directed to the use of isolated hepatoblasts for the treatment of dysfunction and for the production of artificial livers.

BRIEF DESCRIPTION OF THE DRAWINGS

The above brief description, as well as further objects and features of the present invention, will be more fully understood by reference to the following detailed description of the presently preferred, albeit

illustrative, embodiments of the present invention when taken in conjunction with the accompanying drawings wherein:

Figure 1 represents cells from day 14 gestation 5 livers stained for monoclonal antibodies 374.3 followed by FITC and PE-labeled antibodies. Panel A is a two color density plot showing 5 populations designated R1-5 in an ungated sample. and R2 are cell populations positive for OX-43, while 10 R3-5 are negative for this marker. Panel B biparametric dot plot of FL2 versus SSC showing the gating parameters $0X-43^{+}$ used to separate OX-43 cells. The insert shows the negative control. Panel C is a 3D plot of FLl versus FL2 of OX-43 cells 15 showing three distinct cell populations, R3-5;

Figure 2, panel A is a Western blot of total protein from sorted cells showing the presence of albumin containing cells exclusively in the OX-43 population. Panels B and C show indirect immunofluorescence for AFP on OX-43 (B) and OX-43 (C) cells;

Figure 3 represents cells from R3-5 which were sorted after gating out all OX-43⁺ cells and total RNA prepared by the guanidinium isothiocyanate method. The Northern blot demonstrates expression of albumin in R4, while serglycin is expressed by R3 cells;

Figure 4 represents cells which were gated to separate populations positive and negative to OX-43 and then further separated to 5 populations based on their fluorescence on biparametric density plots of FL1 versus FL2. Freshly sorted and cytospun cells were stained for morphology by Diff-Quik staining kit. Original magnification - 100X;

Figure 5 represents a population highly 35 enriched for fetal liver parenchymal cells which was obtained by FACS (R4 cells after exclusion of all OX-43⁻) and 5 x 10⁴ cells/cm² plated on type I

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collagen coated dishes in a serum free, hormonally defined medium. Panel A is a phase micrograph showing a typical epithelial colony and very few mesenchymal cells after 4 days in culture (original magnification - 50%). indirect in situ immunofluorescence 5 Panel B is an showing incorporation of BrdU in the nuclei of about 25% of the cultured parenchymal cells after 24 hours in culture (original magnification - 50X. Panel C is a phase micrograph of panel B;

flow diagram ο£ represents 6 Figure 10 enrichment utilizing a method of the hepatoblast invention;

Figure 7 panel A represents phase contrast microscopy and panel B represents immunofluorescence for 15 AFP of hepatoblasts at gestation day 15. AFP positive cells ranged in morphology from small cells with oval nuclei and scant cytoplasm that were only slightly larger than the hemopoietic cells to cells with larger Negative controls amounts of vacuolated cytoplasm. consisted of cells stained with rabbit IgG as a primary 20 antibody;

Figure 8 represents Northern blot analysis of (5 µg/lane) from freshly isolated fetal RNA liver cells before and after panning and hybridized with encoding α -fetoprotein and albumin. shows freshly isolated fetal liver cells. Lane 2 shows cell preparation after panning 2X with anti-rat RBC Also shown are blots for 18S, used as an antibody. internal control for total RNA loading;

Figure 9 represents biparametric analysis of fetal rat liver cells presented as side scatter (SSC), a complexity, versus of cytoplasmic measure fluorescence for OX-43 and OX-44. Panel Α unstained cells; panel B shows the cells immediately 35 following isolation (original suspension); and panel C shows the cells after final panning. The vast majority of the cells immediately after isolation were agranular and positive for the markers (R1 cell population). With enrichment, the population of granular cells (SSC >50 A.U.) which were negative for the OX43/OX44 markers (R3 cell population) increased. Sorting for this population revealed that 75% were positive for AFP. The demarcation between positive and negative is higher for the granular than the agranular populations due to greater autofluorescence of the granular cells;

Figure 10 represents day 15 gestation cells enriched for hepatoblasts by panning out RBCs cultured for 5 days on type IV collagen in serum-free hormonally defined medium. The cells exhibited typical epithelial morphology including formation of bile canaliculi. Surrounding epithelial cells are fibroblast-like cells.

15 Bar = 25μ ; and

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Figure 11 represents small epithelial islands showing positive staining for albumin by in situ immunofluorescence after 16 days in culture. The fibroblast-like cells surrounding them are negative for the presence of albumin. Bar = 100μ .

DETAILED DESCRIPTION OF THE INVENTION

This invention relates to isolated hepatoblasts and to methods of isolating hepatoblasts from dissociated liver cells utilizing panning techniques and fluorescence activated cell sorting. The isolated hepatoblasts of the invention can be used to treat liver dysfunction, to produce artificial livers, in the study of liver functions, in gene therapy, in drug testing and in vaccine production.

Livers are dissociated by enzymatic digestion, avoiding enzymes such as pronase that adversely affect hepatoblasts, and then kept in solutions which are chilled and which contain chelating agents such as EGTA, which results in cells that can be sustained as single cells. Dissociated liver cells are then panned with antibodies to greatly reduce the numbers of contaminating cell types (hemopoietic cells, including

red blood cells, endothelial cells and other mesenchymal cells in embryonic and neonatal liver, and mature liver cells, hepatocytes, bile duct cells, endothelial cells and other mesenchymal cells in adult liver). Panning alone, although rapid, is inefficient and does not yield very pure cell populations. However, it is used to rapidly reduce the number of non-hepatoblast cells. The cells that do not adhere to the panning dishes are then segregated by fluorescence activated cell sorting, a technology with very high accuracy and efficiency. The combination of the rapid panning methodology with the accuracy of the fluorescence activated cell sorting results in highly purified cell populations with good viability.

In embryonic and neonatal livers, the contaminant cell types reduced through panning protocols are erythroid, myeloid and other hemopoietic cell types and endothelia (mesenchymal cell types). The panning steps lead to a cell population enriched for immature hepatic cell types. In adult livers, the contaminant cell types are mature hepatocytes, bile duct cells, endothelia and some hemopoietic cell populations.

Panned cells are also sorted for multiple subcategories markers that distinguish distinct cell populations. The precursor 25 hepatic identified are (a) the extent of granularity as measured by side scatter on fluorescence activated cell sorting, populations more cell immature wherein more agranular, and increasing granularity correlates with 30 increasing maturity; (b) the extent of autofluorescence, correlates wherein increasing autofluorescence with increasing maturity; and/or (c) the expression of a hepatic cell marker (such as the oval cell marker OC.3, which is detected by monoclonal antibody 374.3).

Liver cells which do not express hemopoietic or endothelial cell antigens recognized by monoclonal antibodies OX-43 and/or OX-44 (which recognize myeloid

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cells and endothelia) and which do not express antigens recognized by a monoclonal antibody to an erythroid antigen comprise the hepatoblasts of the invention. The hepatoblasts of the invention include three categories of immature liver cells:

- (1) More granular cells, which are OC.3⁺, are committed bile duct precursors. These cells are also AFP⁺, albumin⁺ and CK 19⁺.
- (2) More granular cells, which are OC.3, are committed hepatocyte precursors.

 These cells are also AFP, albumin+++, and CK 19.
- (3) Agranular cells, which are OC.3⁺, are very immature hepatic precursors. These cells are also AFP⁺⁺⁺, albumin⁺ and CK 19⁻.

This invention is further directed to the use the methods isolated by hepatoblasts The isolated hepatoblasts of the invention 20 invention. can be used for to treat liver dysfunction. example, hepatoblasts can be injected into the body, such as into the liver or into an ectopic site. Whole requires liver transplantation, which costly dangerous major surgery, can be replaced by a minor surgical procedure which introduces hepatoblasts in vivo either into the liver via the portal vein or at an addition, ectopic site such as the In spleen. hepatoblasts, can be used in bioreactors or in culture livers. artificial form to apparatus hepatoblasts can be used in gene therapy, drug testing, vaccine production and any research, commercial or therapeutic purpose which requires liver cells varying extents of maturity.

<u>Example I</u>

Fischer 344 rats with known durations of pregnancy were obtained from Harlan Sprague Dawley, Inc.

(Indianapolis, IN) and maintained in the animal facility of the Albert Einstein College of Medicine, Bronx, NY on a standard rat chow diet with 12 hour light cycles. By convention, the first day of gestation is defined as day 0. Use of animals was in accordance with the NIH Policy on the care and use of laboratory animals and was approved by the Animal Care and Use Committee of the Albert Einstein College of Medicine.

In order to isolate fetal liver cells, pregnant 10 rats at the fourteenth day of gestation were euthanized with ether and the embryos were removed intact and placed into ice cold CA+2-free Hank's Balanced Salt Solution containing 0.04% DNAse, 0.8 mM MgCl2, HEPES, pH 7.3 (HBSS). Livers were then dissected from 15 the fetuses and placed into fresh ice-cold HBSS. After tissues were collected and non-hepatic tissue removed, HBSS-5 mM EGTA was added to a final EGTA concentration of 1 mM. The livers were moved to a 50 ml conical centrifuge tube by pipette, gently triturated 6 20 to 8 times to partially disaggregate the tissue and then centrifuged at 400 g for 5 minutes at 4°C. subsequent centrifugation steps were performed at the The supernatant was removed and the same settings. pellet of cells and tissue was resuspended in 50 ml 0.6% 25 Collagenase D (Boehringer Mannheim, Indianapolis, IN) in HBSS containing 1 mM CaCl2, gently triturated and then stirred at 37°C for 15 minutes in an Erlenmeyer flask. The dispersed cells were pooled, suspended in HBSS containing 1 mM EGTA and filtered through a 46 μm 30 tissue collector (Bellco Glass, Inc., Vineland, NY). The cell suspension was centrifuged and the cells were resuspended in HBSS supplemented with MEM amino acids, MEM vitamins, MEM non-essential amino acids, transferrin $(10 \mu g/m1),$ (10 μg/ml), iron-saturated 35 free fatty acids (7.6 mEq/L, as described by Chessebeuf Nu-Chek-Prep, Elysian, MN), al., 1984, elements, albumin (0.1%, fraction V, fatty acid free,

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Miles Inc., Kankakee, IL), myo-inositol (0.5 mM) gentamicin (10 μg/ml, Gibco BRL, Grand Island, NY) (HBSS-MEM). Cell number and viability were determined by hemacytometer and trypan blue exclusion.

In order to remove erythroid cells, panning dishes were prepared according to the procedure of Wysocki and Sato (1978) using a rabbit anti-rat RBC IgG Antibodies Gilbertsville, PA). Inc., (Rockland (0.5 mg/dish) diluted in 9 ml of 0.05 M Tris pH 9.5 were poured on 100 mm² bacteriological polystyrene petri dishes (Falcon, Lincoln Park, NJ). The dishes were swirled to evenly coat the surface and incubated at room temperature for 40 minutes. The coated dishes were washed four times with PBS and once with HBSS containing 0.1% BSA prior to use. 15

cell suspension milliliters of the Three containing up to 3 x 10^7 cells were incubated at 4°C for 10 minutes in the dishes coated with the rabbit anti-rat RBC IgG. The non-adherent cells were removed 20 by aspiration and the plates were washed three times with HBSS-0.1%BSA-0.2 mM EGTA and centrifuged. pellet was resuspended in HBSS-MEM and RBC panning was repeated. Following the second RBC panning cell number and viability were determined again.

The cells recovered after RBC panning were then suspension by incubating with mouse labeled in monoclonal antibody OX-43 (1/200=15 μg/ml, 276, MCA Indianapolis, IN) for Science, Bioproducts monoclonal antibody 374.3 (1/500-1/750, a gift of R. Faris and D. Hixon, Brown University, Providence, RI) 30 simultaneously at 4°C for 40 minutes. OX-43 recognizes an antigen on endothelial cells, a subpopulation of erythroid cells (see macrophages and Immunology, Vol. 42, pp. 593-600 (1981) and Robinson et al., Immunology, Vol. 57, pp. 231-237 (1986)) and 374.3 recognizes oval cells, bile duct cells and hemopoietic (see Hixon et al., Pathology: Liver cells

Carcinogenesis, pp. 65-77 (1990)). Second antibodies were PE-conjugated anti-mouse IgG, heavy chain specific (Southern Biotechnology Inc., AL) and FITC-conjugated anti-mouse IgM, heavy chain specific (Sigma Chemical Co., St. Louis, MO). Negative controls included cells without label and cells labeled with mouse isotype controls.

Cells before and after sorting were maintained After completion of the 4°C and in HBSS-MEM. iodiđe propidium labeling, 10 antibody concentration of 10 μ g/ml was added to each of the sample tubes. Fluorescence Activated Cell Sorting was performed with a Becton Dickinson FACSTAR plus Jose, CA) using a 4W argon laser with 60 mW of power and Fluorescent emission at 488 nm 100 µm nozzle. 15 a excitation was collected after passing 530/30 nm band pass filter for FITC and 585/42 nm for PE. Fluorescence measurements were performed using logarithmic amplification on biparametric plots of FL1 20 (FITC) vs FL2 (PE). Cells were considered positive when fluorescence was greater than 95% of the negative control cells.

For measurement of physical characteristics of the cells, FACSTAR plus parameters were FSC gain 8 and SSC gain 8. These settings allowed all cells to be visualized on scale. HBSS was utilized as sheath fluid. For analysis, a minimum of 10,000 events were measured. List mode data were acquired and analyzed using LysisII software. Dead cells were gated out using propidium iodide fluorescence histograms on unlabeled cells.

For determination of positivity to a single antibody dot plots of fluorescence versus side scatter were used. Density plots FL1 versus FL2 were used to select populations with respect to expression of both antigens. A sort enhancement module was utilized for

non-rectangular gating and use of multiparametric gating to select populations of interest.

Shorted cells from day fourteen of gestation all populations were plated in a serum-free, 5 hormonally-defined medium with αMEM as the medium to which the following components were added: EGF (0.01 μg/ml, (10 μg/ml); insulin hormone growth NY); Placid, Lake Biotechnology, (10 μ U/ml); prolactin (20 mU/ml); Triiodothyronine dexamethasone $(10^{-7} \text{ M});$ saturated iron $10 (10^{-7} M);$ transferrin (10 μ g/ml); folinic acid (10⁻⁸ M, BRL, Grand Island, NY), free fatty acid mixture (7.6 as described by Chessebeuf et ai., Nu-Chek-Prep, Elysian, MN); putrescine (0.02 μg/ml); 15 hypoxanthine (0.24 μ g/ml); thymidine (0.07 μ g/ml); bovine albumin (0.1%, fraction V, fatty acid free, Miles IL); trace elements; CuSO₄•5H₂O Kankakee, (0.0000025 mg/l), $FeSO_4 - 7H_2O$ (0.8 mg/l), $MnSO_4 - 7H_2O$ $(0.0000024 \text{ mg/1}), \qquad (NH_4)_6 MO_7 O_2 4^{\bullet} H_2 O \qquad (0.0012 \text{ mg/1}),$ 20 NiCl₂·6H₂O (0.000012 mg/1), NH₄VO₃ (0.000058 mg/1), $\mathrm{H_{2}SeO_{3}}$ (0.00039 mg/l); Hepes (31 mM) and Gentamicin NY). Reagents BRL, Grand Island, (10 μg/ml, Gibco were supplied by Sigma Chemical Company, St. Louis, MO, unless otherwise specified. The trace element mix was a 25 gift from Dr. I. Lemishka, Princeton University, NJ.

Culture dishes as well as cytospins of various cell suspensions were fixed with ice-cold ethanol or acetone. After blocking with PBS containing 1% BSA for 30 minutes at room temperature, the fixed cells were immunofluorescence using by indirect 30 studied following primary antibodies: polyclonal rabbit-anti-rat Biochemical Corporation, States (United albumin Cleveland, OH), rabbit-anti-mouse AFP antiserum (ICN monoclonal Mesa, CA), Costa In., Biomedical, 35 mouse-anti-human cytokeratin 19 (Amersham Life Science,

Arlington Heights, IL), polyclonal rabbit-anti-human IGF II receptor (a gift of Dr. Michael Czech, University of Worchester, MA), mouse monoclonal anti-rat-Thy-1 (OX-7, Bioproducts for Science, Indianapolis, IN), monoclonal 5 mouse-anti-desmin (Boehringer Mannheim, Indianapolis, IN), and 258.26, a monoclonal mouse-anti-rat antibody identifying postnatal hepatocytes as well as some fetal liver parenchymal cells (a gift of Drs. R. Faris and D. Second antibodies RI). Brown University, Hixon, 10 included species specific Rhodamine conjugated antibodies corresponding to the primary antibodies. controls consisted of cells stained with mouse or rabbit IgG or mouse isotype controls. Freshly isolated adult hepatocytes were used as positive controls for albumin Gamma-glutamyltranspeptidase (GGT) 15 staining. assayed by immunochemistry on ethanol fixed cells using the method described by Rutenberg et al., J. Hist. Cyt., Vol. 17, pp. 517-526 (1969).

In order to perform Northern blot analysis for 20 the presence of specific mRNA, total RNA was extracted from sorted cells using the guanidinium isothiocyanate method, as described by Chomcznyski et al., Anal. Biochem., Vol. 162, pp. 156-159 (1987)). RNA samples were resolved by electrophoresis through 1% agarose 25 formaldehyde gels in 3-(N-morpholino)-propanesulfonic acid buffer (see Maniatis et al., Molecular Cloning: A <u>Laboratory Manual</u>, pp. 191-193 (1982)). The RNA was then transferred to Gene Screen (New England Nuclear, Boston, MA), and the filters were prehybridized and hybridized with the appropriate probes. The cDNA clones complementary to specific mRNAs were radioactively labeled by primer extension with 32P dCTP as described by Feinberg et al., Anal. Biochem., Vol. 137, pp. The cDNAs used in hybridization were 266-267 (1984). 35 rat albumin (a gift of Dr. Zern, Jefferson University, α-fetoprotein, mouse and Philadelphia, PA),

(Dr. Tighlman, Princeton, NJ), GGT (obtained from Dr. M. Manson, MRC Medical Research Council, Surrey, UK) and Autoradiograms were scanned with a Quantimat 920; Manufacturer's Cambridge (Model densitometer The data for each of the genes was 5 Instrument). normalized to that for the common gene 18S (J. Darnell, Rockefeller University, New York, NY).

In order to perform Western blot analysis, total protein samples from various sorted cells were 10 loaded on a 10% polyacrylamide minigel. Loading was normalized for equal cell numbers, 100,000 cells per Electrophoresis followed by electroblotting to nitrocellulose membranes (Schleicher and Schuell, Keene, NH) was performed. The blots were blocked overnight in 15 2% dry milk solution at 4°C and assayed for albumin using a rabbit-anti-rat albumin antiserum diluted 1:800 in the blocking solution for 1 hour at room temperature, incubation with hour one by followed horseradish-peroxidase- conjugated anti-rabbit IqG 20 (Amersham Life Science, Arlington Heights, IL) diluted 1:50 in blocking solution. Detection was achieved by incubation of blots with ECL-chemiluminescence kit reagents (Amersham Life Science, ARlington Heights, IL) for 1 minute and subsequent autoradiography.

Forty-eight well plates were coated with type I collagen extracted from rat tail tendon as described by Reid, Methods in Molecular Biology, The Humana Press, Inc., Vol. 5, pp. 237-276 (1990). Sorted cells at densities between 50,000 to 100,000 cells/cm² were plated per well. Following an overnight attachment period, the medium with the non-adhering cells was gently removed and replaced by fresh medium. A complete medium change was performed every 24 hours. The cells were cultured at 37°C in a fully humidified atmosphere 35 containing 5% ${
m CO}_2$ and were observed daily. After four days in culture, cells were fixed with ice-cold ethanol and stained in situ by Immunofluorescence for albumin,

AFP, CK 19 and IGF II receptor and by immunochemistry for GGT, as described below.

Livers from fourteenth day gestation embryos isolated by the EGTA-collagenase digestion yielded 5 single cell suspensions and a negligible number of cell Cellular viability was greater than 95% as aggregates. determined by exclusion of trypan blue. Cell yield was $2.62 + 0.31 \times 10^6$ cells per liver. The original cell suspension was subjected to two steps of immunoadherence anti-rat RBC ("panning") using rabbit IgG polystyrene dishes. Cellular recovery after completion of two panning steps was 51% (\pm 8%), but varied somewhat with different lots of antibodies.

The cells recovered after RBC-panning were stained in suspension with a mixture of two antibodies: 15 an antibody raised against "oval cells" (monoclonal antibody 374.3) and a commercially available antibody recognize endothelial, as well as some erythroid and myeloid cells in the rat (monoclonal 20 antibody OX-43). Following incubation with the proper FITC and PE labeled second antibodies, cells were analyzed for their fluorescence patterns. As shown in Figure 1, panel A, when fluorescence intensities for both antigens were plotted against each other, five 25 distinct populations, referred to as R1 through R5, were observed. With minor variations in the percentage of each population, the distribution of cells to form the five populations was extremely reproducible. differences could be explained by variations in the percent recovery of cells after RBC panning. 30

of cells Initial analyses sorted by immunofluorescence revealed the presence of albumin and AFP positive cells in one of the OX-43 positive cell populations (R2). These larger and more complex cells 35 comprised approximately 5-10% of cells in this gate. However, when freshly sorted R2 cells were viewed under the epi-fluorescent microscope, these larger

appeared to be negative for OX-43 (no PE labeling). parenchymal cells in the liver have a significant degree of autofluorescence, which increases with maturation of the liver, in parallel to the increase in cellular complexity, as measured by the side scatter parameter on the FACS. It was therefore postulated that it is due to this phenomenon that some parenchymal cells appear in the region of the OX-43-positive cells, although not To pursue this hypothesis, expressing the antigen. positivity to OX-43 was determined accurately on side scatter (cellular granularity) versus PE fluorescence, as measured on the FL2 scale (Figure 1, panel B), and OX-43-positive and negative cells were sorted and To determine the accuracy of the sorts, characterized. cells ο£ sorted the acquisitions post-sort performed using the same instrument settings. post-sort purity (i.e., percentage of cells from a shorted population that appeared in the same region when analyzed again after the sort) was >90%.

Sorted cells from both OX-43 positive and 20 negative gates were assayed for expression of liver specific genes by Western blot analysis and by indirect As shown in Figure 2, panel A, immunofluorescence. albumin in of the minimal amount there was а fraction, detected by Western cell OX-43-positive blotting, as compared with the OX-43-negative cells. No AFP positive cells could be shown by indirect immunosorted OX-43-positive on cytospins of fluorescence as opposed to 30% of OX-43-negative cells expressing the fetal liver marker (see Figure 2, panels 30 B and C). It was concluded that at day 14 of gestation, all fetal liver parenchymal cells are OX-43-negative. achieve "cleaner" in order to Therefore, OX-43-positive and negative cells were separated on a SSC versus FL2 plot and studied separately.

When OX-43 positive cells were electronically gated out and the remaining cells viewed on a FL1 versus

FL2 plot, three distinct populations were readily detected (see Figure 1, panel C), corresponding to R3-5 in the ungated cell suspension. All of the cells in R3 were 374.3-positive whereas 30% of the cells in R4 were positive for that marker. R5 cells did not express OC.3. Expression of various liver-specific and other genes was studied on sorted cells from R3-5. The results are summarized in Table 1, below.

TABLE 1

Characterization of sorted cells by immunofluorescence and by histochemistry

		R1	R2	R3	R4	R5
	Albumin	neg	neg	1% pos	75-80% pos	neg
	AFP	neg	neg	2% pos	70% pos	neg
L5	GGT	neg	neg	1% pos	75%	neg
	IGF-IIr	20%	1%	2%	85%	neg
	CK 19	neg	neg	2-3%	neg	neg
	Desmin	<1% +	1-2% +++	neg	neg	<1% +
	258.26	neg	neg	neg	neg	neg
20	Thy-1	2%	10%	75%	10%	5%

About 2-3% of R3 cells (less than 0.2% of the total ungated cell suspension) were intensely stained for albumin and AFP. They also expressed GGT and CK 19, markers of the bile duct lineage. However the majority of the cells appeared to be small, blast-like cells, and did not express liver specific genes but expressed Thy-1 and hemopoietic markers such as classical serglycin (see Table 1 and Figures 3 and 4). Most of the liver parenchymal cells were found in the R4 gate 30 (see Table 1 and Figure 3). The vast majority of the cells expressed albumin, AFP and GGT, all markers of

fetal liver parenchyma. No hemopoietic or fat storing cell markers were detected in that gate. The cell population designated R5 is a heterogeneous one (see Figure 4), comprising mainly two cell types: (1) cells that morphologically appear to be normoblasts; and (2) simple small cells that did not express parenchymal liver genes. The ratio between these two cell types varied somewhat and was dependent on the efficiency of the RBC panning.

When all of the OX-43 negative cells were gated 10 two distinct populations were observed As expected, no parenchymal liver markers FL1/FL2 plot. were detected in these cells. A few of R2 cells intensely stained with the antibody against desmin, an intermediate filament usually expressed in fat storing 15 cells. Morphologically, most of R2 cells appeared to be early erythroid precursors (see Figure 4), while 10% of R1 gate were the expressed Thy-1. In morphologically distinct cell types (see Figure 4). majority were small, blast-like and did not express any 20 The others, about 20% of the of the markers tested. cells in this gate, were larger cells with a pale cytoplasm and expressed the receptor for IGF-II. Very few cells from Rl stained for Thy-1.

Sorted cells from all 5 populations were cultured for 4 days to determine <u>in vitro</u> fates. When plated at high density under the conditions described, R4 cells yielded clusters of epithelial cells surrounded

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by very few scattered stromal cells (see Figure 5A and Table 2 below).

TABLE 2

Characterization of R4 cells

after 4 days in culture

	Marker	Epithelial Cells	Stromal Cells
	Albumin	+	neg
	AFP	±	neg
	GGT	++	neg
.0	CK 19	+(30%)	neg
	258.26	neg	neg
	IGF IIr	+ (perinuclear staining)	+ (perinuclear staining

Cell division was clearly evident both in the epithelial as well as the stromal components of the culture. On the second day of the culture 25±5% of the epithelial cells showed incorporation bromo-deoxy-uridine (BrdU) following one hour incubation with a medium containing BrdU (see Figure 5A and B). When RBC-panned but not sorted day 14 gestation 20 cells were plated under similar conditions, survived for at least 10 days (data now shown). of sorted R4 cells However, cultures deteriorated The epithelial cells lost their classical quickly. polygonal shape and elongated, similarly to what is seen 25 in primary cultures of adult hepatocyte in the presence Moreover, when stained in situ for albumin, AFP and GGT, cultured R4 cells exhibited a gradual liver-specific decline in these genes, RBC-panned day 14 gestation cells maintained their gene expression under similar conditions (data not shown). IGF-II receptor remained clearly detected in the golgi of the cultured epithelial as well as the stromal

cells. About 30% of the cultured R4 cells showed staining for CK 19, a cytokeratin present in bile duct cells and not in adult hepatocytes.

When cells from all other four populations were 5 plated under the same conditions, only few scattered fibroblast-like cells (but not epithelial colonies) were observed. Despite the liver-parenchymal characteristics of some R3 cells, epithelial colonies from these cells could not be obtained under similar plating conditions. 10 This may have been due to low density of the epithelial cells aggregated These gate. this in suspension, survived for about 48 hours and then died. Coating the dishes with type I or type IV collagen, fibronectin or laminin alone or in combination did not 15 improve attachment or survival of these cells (data now shown).

Example II

Fisher 344 rats with known durations of pregnancy were obtained from Harlan Sprague Dawley, Inc.

(Indianapolis, IN) and maintained in the animal facility of the Albert Einstein College of Medicine, Bronx, NY on a standard rat chow diet with 12 hour light cycles. By convention, the first day of gestation is defined as day O. Use of animals was in accordance with the NIH Policy on the care and use of laboratory animals and was approved by the Animal Care and use Committee of the Albert Einstein College of Medicine.

Pregnant rats at the fifteenth day of gestation were euthanized with ether, and the embryos were delivered. Livers were then dissected from the fetuses, weighed, placed into ice-cold, Ca⁺²-free Hank's Balanced Saline Solution containing 0.8 mM MgCl₂, 20 mM HEPES, pH 7.3 (HBSS), and gently agitated at room temperature for 1 minute. After removal of non-hepatic tissue, livers were gently triturated and then stirred at 37°C for 10 to 15 minutes in an Erlenmeyer flask with 0.6% type IV collagenase (Sigma Chemical Co., Lot

11H6830, St. Louis, MO) in HBSS containing 1 mM CaCl, and 0.06% DNAse I (Boehringer Mannheim, Indianapolis, minute intervals, tissue fragments were At 5 The supernatant sediment at lq. allowed to recovered and fresh collagenase solution added. The **HBSS** suspended pooled, were dispersed cells containing 5 mM EGTA and filtered through Inc., Vineland, tissue collector (Bellco Glass, under lg. The resultant cell suspension was centrifuged at 4°C for 5 minutes under 450g. The cell pellet was 10 resuspended in HBSS containing 0.2 mM EGTA and 0.5% BSA (HBSS-EGTA-0.5% BSA), and the cell number was estimated with a Coulter Counter (Coulter Electronics, Hialeah, FL). Cell viability was assessed by exclusion of 0.04% trypan blue, and an aliquot of the suspension was centrifuged in a tared microfuge tube at 450g for 5 minutes.

immunoadhere hemopoietic order In onto antibody-coated polystyrene endothelial cells dishes, panning dishes were prepared according to the 20 The antibodies employed procedure of Wysocki and Sato. IgG (Inter-cell RBC anti-rat rabbit included Technologies, Inc., Hopewell, NJ) and goat Ig3 directed towards mouse whole IgG molecule (M-3014, Sigma, St. 25 Louis, MO). Antibodies (0.5 mg/dish) diluted in 10 ml 9.5 were poured on рΗ 0.05 M Tris dishes (Falcon, petri polystyrene bacteriological NJ) to evenly coat the surface Lincoln Park, incubated at room temperature for 40 minutes. coated dishes were washed four times with PBS and once with HBSS containing 0.1% BSA prior to use.

Three milliliters of the cell suspension containing up to 3 x 10⁷ cells were incubated at 4°C for 10 minutes in the dishes coated with the rabbit anti-rat RBC IgG. The supernatant containing non-adherent cells was removed by gentle aspiration while tilting and swirling, combined with three washes

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of 7 ml HBSS-EGTA-0.1% BSA, and centrifuged at 4°C for 5 minutes under 450g. Cells from two dishes were pooled and repanned with a fresh dish coated with rabbit anti-rat RBC IgG. The non-adherent cells were then 5 removed as above and resuspended with HBSS-EGTA-0.5% BSA $1 \times 10^{7}/\text{ml}$. The of concentration to hepatoblasts were then incubated simultaneously at 4°C for 40 minutes with mouse monoclonal antibody OX-43 Indianapolis, Serotec, MCA276, (15 μg/ml, monoclonal antibody OX-44 (18 μ g/ml, MCA371, Serotec, 10 OX-43 recognizes an antigen on Indianapolis, IN). macrophages, endothelial cells and red blood cells, and OX-44 recognizes the membrane-glycoprotein CD53 that is present on all rat myeloid cells as well as peripheral lymphoid cells, and is related to the human leukocyte antigen CD37. After washing to remove excess antibody, cells were panned at 4°C for 10 minutes in a dish coated the goat anti-mouse whole IgG antibody, non-adherent cells were removed as described above.

Next, cytospins of the various cell suspensions 20 were fixed with either ice-cold ethanol or alcohol, acetone and carbowax 1540 (Fix-Rite, Richard-Allan Medical Industries, Richland, MI). After blocking, the indirect immunostained by were fixed cells 25 immunofluorescence or the biotin/streptavidin method using B-galactosidase (BioGenex, San Ramon, CA) with rabbit anti-rat albumin IgG (USB Corp., Cleveland, OH) (ICN antiserum anti-mouse AFP rabbit or as primary antibodies. ImmunoBiologicals, Lisle, IL) 30 Negative controls consisted of cells stained with the Positive controls antibodies omitted. albumin staining were done with freshly isolated adult hepatocytes.

In order to perform Northern blot analysis, total RNA was extracted from the cells before and after panning and from the cells adherent to the panning dishes using the guanidinium isothiocyanate method.

samples were resolved by electrophoresis through 1% gels formaldehyde agarose buffer, then 3-(N-morpholino)-propanesulfonic acid transferred to Gene Screen (New England Nuclear, Boston, MA), which was prehybridized, and then hybridized with the appropriate probes. The cDNA clones complementary to specific mRNAs were radioactively labeled by primer extension with 32 P dCTP. The cDNAs used were rat 18S (J. Darnell, mouse and mouse AFP albumin, Autoradiograms University, NY). 10 Rockefeller scanned with a Quantimat densitometer (Model Manufacturer's Cambridge Instrument). The data for each of the genes was normalized to that for the common gene 18S.

To perform FACS analysis sorting and 15 hemopoietic and endothelial cell markers at day 15 stages of various suspensions at cell gestation, enrichment were analyzed by flow cytometry in the FACS facility of the Albert Einstein College of Medicine, Bronx, NY. Cells were resuspended to 1×10^7 cell/ml and incubated at 4°C for 40 minutes with OX-43 with and without OX-44, followed by FITC-conjugated anti-mouse IgG (heavy chain specific, Southern Biotech, Birmingham, AL) at 4°C for 40 minutes. Cells stained only with negative anti-mouse IgG served as FITC-conjugated controls.

Flow cytometric analysis was performed on a Becton-Dickinson FACScan (San Jose, CA) with a 15mW air-cooled argon laser. Cell sorting was performed with 30 a Becton Dickinson FACSTAR plus (San Jose, CA) using a 4W argon laser with 60mW of power and 100 μm nozzle. fluorescent emission 488 nm at instances both excitation was collected after through passing Fluorescence filter for FITC. pass 530/30 nm band logarithmic using performed were 35 measurements Cells were considered positive when amplification. fluorescence was greater than 95% of the negative

physical measurement of For control cells. characteristics of the cells, the detector value was E-1 for forward scatter (FSC) with mid-range amplification. For side scatter (SSC) the detector value was mid-range 5 with an amplification of 1. Equivalent FACSTAR plus parameters were FSC gain 4 and SSC gain 8. settings allowed all cells to be visualized on scale. using were performed gating SSC and into 256 parameters both dividing amplification, arbitrary units (A.U.). For analysis, at least 10,000 10 events were measured. List mode data were acquired and analyzed using LysisII software. Cells before and after sorting were maintained at 4°C and in HBSS supplemented with insulin, transferrin, free fatty acids, detailed gentamicin as 15 elements, albumin, and supplements added to the HDM.

Next, multiparametric flow cytometric analysis of hemopoietic and endothelial markers was performed with respect to the oval cell antigen OC.3. Isolated 20 cells were labeled with a combination of OX-43 and OX-44 (mouse IgGs) and monoclonal antibody 374.3 (mouse IgM, Hixson and Faris, Brown University, Providence, followed by FITC-conjugated goat anti-mouse IGG (heavy Birmingham, Biotech, specific, So chain PE-conjugated goat anti-mouse IgM (heavy chain specific, 25 Cells stained only So Biotech, Birmingham, AL). PE-conjugated IqG and anti-mouse FITC-conjugated anti-mouse IgM served as negative controls. Cells were evaluated both for extent of fluorescence for one of the probes and by side scatter, a measure of cellular complexity (extent of cytoplasmic organelles).

Cells from day 15 gestation livers were panned antibody, the and cell blood red rat against epithelial-enriched cell suspension was plated in a serum-free hormonally defined medium with αMEM as the basal medium to which the following components were insulin (10 μg/ml); EGF (0.01 μg/ml, Upstate added:

Biotechnology, Lake Placid, NY); growth hormone (10 μ U/ml); prolactin (20 mU/ml); glucagon Triiodothyronine (10⁻⁷M); dexamethasone (10 μg/ml); saturated transferrin (10 μg/ml); $(10^{-7}M)$; iron folinic acid (10⁻⁸M, Gibco BRL, Grand Island, NY), free fatty acid mixture (0.76 mEq/1, a modification of the method described by Chessebeuf, Nu Check-Prep, putrescine (0.02 µg/ml); hypoxanthine Elysian MN); (0.24 μ g/ml); thymidine (0.07 μ g/ml); bovine albumin 10 (0.1%, fraction V, fatty acid free, Miles Inc., Kankakee, IL); trace elements: $CuSO_4 - 5H_2O$ (0.0000025 mg/l), $FeSO_4 \cdot 7H_2O$ (0.8 mg/1), $MnSO_4 \cdot 7H_2O$ (0.0000024 mg/1), (0.0012 mg/1), NiCl₂-6H₂O $(NH_4)_6MO_7O_{24} \cdot H_2O$ $(0.000012 \text{ mg/l}), NH_4 VO_3$ (0.000058 mg/l),15 (0.00039 mg/l); Hepes (31 mM) and Gentamicin (10 μ g/ml, Gibco BRL, Grand Island, NY). Reagents were supplied by Sigma Chemical Company (St. Louis, MO) unless otherwise specified. The trace element mix was a gift from Dr. I. Lemishka, Princeton University, NJ. 20

Twenty-four well plates were coated with type IV collagen extracted from EHS tumors. Panned cells at densities between 12,500 and 25,000 cells per cm² were plated per well and allowed to attach for four to five hours after which the medium with the non-adhering cells were gently removed and replaced by fresh medium. Cells were cultured at 37°C in a fully humidified atmosphere containing 5% CO₂ and were observed daily for 5 to 16 days. A complete medium change was performed every 48 hours.

At various time points after initiation of the culture, cells were fixed with ice-cold ethanol and stained in situ by immunochemistry or by immunofluorescence for albumin and AFP.

The weight of the liver at the 15th day of 35 gestation was 9.1 ± 1.3 mg. Collagenase treatment

digested the liver completely, and only minimal particulate matter was excluded by the tissue sieve. The number of cells obtained at this step was 1.07 x 10⁷/liver, and the weight of the dissociated cells was 8.6 ± 1.1 mg/liver, 95% of the whole organ weight. The suspension consisted almost entirely of isolated single cells with occasional small aggregates that increased in size and number in the absence of EGTA and at temperatures greater than 4°C. Viability by trypan blue exclusion was greater than 90%.

After each panning, phase contrast microscopy demonstrated that the adherent cells exhibited erythroid morphology. Only rare cells were positive for albumin by immunochemistry. After panning with the rabbit anti-rat red blood cell antibody-coated dishes to 15 remove red blood cells and then with the goat anti-mouse whole molecular IgG antibody-coated dishes to reduce the numbers of OX43/OX44⁺ cells, the non-adherent cells constituted 29 \pm 5% and 16 \pm 4%, respectively, of the 20 cell number of the freshly dispersed fetal liver (original suspension). Panning proved successful for liver tissue at all fetal and early neonatal ages, although the variation in hemopoietic constituents with developmental age resulted in differing degrees of 25 enrichment (data not shown). Also, the efficiency of the RBC panning procedure varied with the antibody lot. With antibodies of poor efficiency for direct panning, however, indirect immunoadherence was successful for the cells labeled in suspension followed by panning with 30 anti-rabbit IgG coated petri dishes.

On phase contrast microscopy following liver dispersion the predominant cell type was a small, red cell consistent in morphology with that of an early erythroid cell. Also present were larger, vacuolated cells. Immunocytochemistry demonstrated that the vast majority of the vacuolated cells as well as occasional smaller, oval-shaped cells were strongly positive for

albumin and AFP (see Figure 7). The proportions of albumin and AFP positive cells at various stages of enrichment are shown in (see Table 3 below and Figure 6).

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TABLE 3

Characteristics of the E15 liver cellular suspension at various stages of enrichment

Markers	Percent of cells positive in the Original Suspension	Percent of cells positive after RBC Panning	Percent of cells positive after IgG Panning
Albumin ¹	3.2 ± 1.3	9.5 ± 1.2	14.8 ± 3.6
Alpha-fetoprotein	2.5 ± 0.7	9.8 ± 0.9	14.9 ± 2.5
MoAb OX-432	76.6 ± 5.8	70.5 ± 6.1	ND
MoAb OX-43/44 ²	87.9 ± 2.5	80.4 ± 3.9	69.0 ± 10.0
% cells remaining of original suspension	100	29 ± 5	16 ± 4

1 Immunocytochemistry with the biotin/streptavidin method using B-galactosidase (BioGenex, San Ramon, CA) with primary antibody omitted as negative control. 2Cells were considered positive when fluorescence was greater than 95% of the negative control cells by FACS analysis. ND = Not done

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Northern blot analysis for liver-specific genes (albumin and AFP) was done on cells before and after panning and is shown in Figure 8. The cells after panning were enriched up to 5-fold for AFP mRNA and 2-fold for albumin mRNA, a finding indicative both of the success of the panning procedures and of the high concentrations of hepatoblasts (as opposed to mature hepatocytes). Negligible levels of albumin and no AFP mRNA were evident in the cells adherent to the panning dishes.

efficiency with which То determine the hemopoietic and endothelial cells were removed, cells at various stages of enrichment were analyzed by flow cytometry for the presence of OX-43 which recognizes macrophages, endothelial cells and red blood cells and for the presence of OX-44 which recognizes myeloid and peripheral lymphoid cells. The results are shown in Figure 6 and in Table 3. The percentage of cells positive for OX-43/OX-44 in the original cell suspension The combination of panning procedures was 87.9 ± 2.5%. and anti-mouse whole anti-rat RBC IqG with of . the antibodies removed 84% cells. Although 69 ± 10.0% of the non-adherent cells were still positive for the OX-43/44 markers, the percentage of hepatoblasts was enriched dramatically (5-fold). Although additional could reduced the $0X - 43/44^{+}$ cell have panning population even further, it was found that the cell numbers had been reduced sufficiently by panning to enable the FAC sorting to complete the process of eliminating the OX-43/44 cells.

When examined by flow cytometry, fetal liver cells constituted a heterogeneous population with respect to FSC, a measure of cell size, and SSC, a measure of cytoplasmic complexity. Cytologically, there was a broad range in cell size (5 to 15 μ by Coulter Counter, data not shown), but cell size was not found to be useful in separating hemopoietic from parenchymal

precursors. Rather, the populations were best segregated using SSC. The definition of granular versus agranular cells was made based on a linear scale for side scatter using biparametric plots of fluorescence versus side scatter. Based on the population profiles, 50 A.U. usually demarcated the agranular from the granular cells.

Using SSC versus fluorescence, the fetal liver populations: three into isolated could be cells 10 agranular cells (the Rl population), which were positive for the endothelial and/or myeloid markers (0X43/0X44), and agranular (R2) and granular (R3) cells negative for the OX43/OX44 markers (see Figure 9). The demarcation and negative was higher for between positive granular than the agranular populations due to greater 15 autofluorescence of the granular cells. Analysis of the sorted FACS populations demonstrated that less than 1% $3.0 \pm 0.7\%$ of the cells in the Rl R2 populations, respectively, were positive for However, 75.1 \pm 4.7% of the granular cells negative for 20 positive for were markers (R3) immunocytochemistry (see Table 4 below).

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FABLE 4

Characteristics of cell fractions on FACS

***	R1	R2	R3
Fluorescence for 276 and/or 3711	positive	negative	negative
Granularity (A.U.) 2	agranular	agranular	granular
% AFP positive ³	< 1%	3.0 ± 0.7%	75.1 ± 4.7%

greater positive when fluorescence was lcells were considered positive when fluorescence than 95% of the negative control cells by FACS analysis. $^{
m lCells}$

250~A.U. demarcated the agranular from the granular cells using FACS parameters of FSC gain 4 and SSC gain 8.

the biotin/streptavidin method using San Ramon, CA) with primary antibody 3Immunocytochemistry with B-galactosidase (BioGenex, omitted as negative control.

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analysis of the Rl cell Double image population, the only one analyzed having OX-43/OX44⁺ cells, indicated extensive overlap of OX-43/44 positive The FACS pattern OC.3 positive cells. OX-43/OX-44 was similar for all gestational ages except for a subtle increase in the Rl (and concomitant the R3 population) with increasing in decrease qestational age due to increasing hepatic erythropoiesis not shown). Analysis of the sorted (data population that was positive for OX-43/44, regardless of expression of OC.3 or of granularity, revealed that morphologically most were hemopoietic precursor cells and were negative for AFP. Of the granular, OX-43/44 cells (the R3 cell population), most of which were oc.3⁺. AFP⁺, approximately 30% were Α small cells (R2 in Table 4) population ο£ that were AFP⁺ $0X43/44^{-}$, and have agranular, not been evaluated for OC.3 expression.

Cell preparations from day 15 gestation enriched by panning for hepatoblasts were plated on type 20 collagen-coated dishes and in the serum-free, hormonally defined medium as described. Within a day after plating, the epithelial cells reaggregated and attached to the matrix as small cell clusters. Plating 25 efficiencies of up to 60% were obtained (data not The cells were organized into islands of shown). parenchymal cells forming close cell-cell typical and bile canaliculi, surrounded contacts non-epithelial, fibroblast-like cells (see Figure 10). culture the 30 After 4-5 days in parenchymal cell gradually overgrown were by the components non-parenchymal cells. However, residual clusters of hepatoblasts remained positive for albumin and AFP for up to 16 days in culture, as assessed by in situ immunochemistry or immunofluorescence (see Figure 11). In a few experiments in which glucagon was omitted from medium, noticeable the culture no morphological

difference was observed, and the cells expressed albumin and AFP when stained in <u>situ</u> by immunofluorescence or immunochemistry (data not shown). This observation is attributed to relative glucagon resistance of the fetal hepatoblasts.

developed have inventors The incorporating panning technologies and multiparametric FAC sorting, which isolate cell populations highly enriched for liver parenchymal cell precursors. methods of this invention have been found by the 10 inventors to be applicable to the isolation of hepatic precursor cells from liver from gestational age day 13 through the early neonatal period. The liver dispersion procedure described yields a population of predominantly single cells with greater than 90% viability, and at 15 gestation day 15, 95% of the whole organ weight is The panning procedures remove up to 84% of recovered. the total cell number, and simultaneously enrich the hepatoblast population by 5-fold. The increase in the 20 parenchymal-specific gene expression of albumin and AFP was illustrated by Northern blot analysis of the cells procedure's the after panning, and and the cells analysis of specificity demonstrated by Similarly, dishes. panning the to adherent enrichment was confirmed by the in vitro data in which there was a dramatic increase in the number of cell colonies expressing albumin and AFP after panning Furthermore, the compared to the original suspension. efficiency after panning was significantly plating higher (up to 60%) compared to previously reported Though the hepatoblasts still values of 6 to 10%. remain a minor population after panning procedures, it is important to consider that the standard <u>in situ</u> protocols yields a population hepatocyte perfusion containing, on average, 37.7% hepatocytes. 35

The advantage of this protocol in comparison with previous methods which involved attachment of

dispersed liver cells to culture dishes, low-speed culture centrifugation, and differential arginine-deficient medium are several-fold. Isolate hepatocytes rapidly lose tissue-specific gene regulation 5 in vitro. As a result, in procedures requiring cell measurement matrix, to attachment parenchymal-specific function, such as protein or mRNA content, might not reflect in vivo levels. Dissociated fetal hepatoblasts also readily form large aggregates temperature-dependent, and calcium 10 via glycoprotein-mediated process. As early as gestation day 14, high levels of a cell membrane protein which is thought to be uvomorulin (E-cadherin) were present on hepatoblasts. This tendency for aggregation explains 15 the ability of low speed differential centrifugation to hepatoblasts, relatively large (E19) enrich especially in the presence of Ca2+ and at temperatures greater than 4°C. To disaggregate the hepatoblasts, mechanical methods including vigorous pipetting and 20 aspiration through a syringe have been employed but found to be insufficient, leading to difficulties with further analyses which require a single cell suspension such as FACS.

The tendency of the cells to aggregate 25 prevented by maintaining the cells at 4°C and by interfering with EGTA. with calcium removing CAM-mediated aggregation. The advantage of maintaining the cells as a single cell suspension is two-fold. First, measurement of parenchymal specific functions can determined on a cellular basis, overcoming the physiologically irrelevant changes in hemopoietic cell population. Second, procedures such as FACS which demand a single cell suspension can be easily performed.

Though gestation day 15 hepatoblasts appear 15 larger than the non-parenchymal cells, side scatter 15 rather than forward scatter on the FACS proved to be a 16 better discriminator in separating the various

populations, presumably because even gestation day 12 hepatoblasts, which contain vacuoles, mitochondria and abundant endoplasmic reticulum, are relatively complex. In addition, side scatter proved a reasonable measure of cellular maturity. In general, hepatoblasts of greater granularity were more mature morphologically and biochemically (data not shown).

Hence, FACS analysis was employed to examine the expression of the oval cell marker, OC.3, which has been proposed to identify liver stem cells. analysis for OC.3 or OX-43/44 multiparametric FACS expression in combination with gating for cells of particular levels of granularity, the inventors were able to subdivide the populations into non-parenchymal stromal cells (hemopoietic, endothelial, and cells) versus parenchymal cell precursors that were AFP⁺. the inventors were able to evaluate the Moreover, OC.3 various antigen in the of the expression At gestation day 15, most agranular, subpopulations. OX43/44 cells proved to be hemopoietic cells, largely erythroid cell populations. Of the granular, 0X43/44 which were predominantly population, approximately 30% of the cells were OC.3 and probably represented bile duct cell precursors, whereas the 25 OC.3 cells were probable hepatocyte precursors. a small percentage of agranular, OX43/44 cells were AFP+.

In comparison to the hemopoietic field, the liver stem cell field is still in its infancy. 30 the ability to isolate specific populations by FACS sorting using these parameters with subsequent in vitro and in vivo fate studies will greatly aid in identifying Furthermore, this technology is the liver stem cell. applicable to the study of all aspects of liver stem 35 cell including the biliary epithelium, biology carcinogenesis, regeneration, aging and tissue-specific gene expression.

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Although the invention herein has been described with reference to particular embodiments, it is to be understood that these embodiments are merely illustrative of various aspects of the invention. Thus, it is to be understood that numerous modifications may be made in the illustrative embodiments and other arrangements may be devised without departing from the spirit and scope of the invention.

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WE CLAIM:

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- A method of isolating hepatoblasts from 1. embryonic or neonatal liver comprising:
 - preparing a single cell suspension of (a) embryonic or neonatal liver cells;
 - said suspension utilizing panning (b) antibodies specific for hemopoietic including red blood cells, cells, other OI endothelial cells mesenchymal cells so remove to as including cells, hemopoietic endothelial cells blood cells, from said other mesenchymal cells suspension; and
 - activated fluorescence performing (c) said utilizing sorting cell remove to antibodies 50 cells, including red hemopoietic including blood red blood cells, and other cells, endothelial cells said cells from mesenchymal performing suspension and fluorescence multiparametric said on cell sorting activated least one suspension utilizing at antibody to a hepatic cell marker, side scatter, forward scatter and/or autofluorescence such that the cells said suspension are in remaining isolated hepatoblasts.
- The method of Claim 1 wherein the antibody 30 specific for hemopoietic cells is a monoclonal antibody.
 - method of Claim 2 wherein The monoclonal antibody is OX-43 and/or OX-44.
- The method of Claim 1 wherein the antibody to a hepatic cell marker is monoclonal antibody 374.3. 35

- 5. The method of Claim 1 wherein said hepatic cell marker is OC.3.
- 6. The method of Claim 1 wherein said single cell suspension contains an agent capable of removing 5 calcium from liver cell surface.
 - 7. The method of Claim 1 wherein said single cell suspension contains EGTA.
- 8. The method of Claim 1 wherein said single cell suspension contains an enzyme capable of 10 dissociating liver cells.
 - 9. The method of Claim 1 wherein said single cell suspension contains collagenase.
 - 10. The method of Claim 1 wherein said single cell suspension is chilled.
- 11. The method of Claim 1 wherein said single cell suspension is at a temperature of between about 2 and 20°C.
 - 12. Hepatoblasts isolated by the method of Claim 1.
- 20 13. A method of isolating hepatoblasts from adult liver comprising:
 - (a) preparing a single cell suspension of adult liver cells;
 - utilizing suspension panning said (b) specific for mature antibodies hepatocytes, mature bile duct cells, mesenchymal endothelial cells and remove to so as cells hepatocytes, mature bile duct cells, and mesenchymal endothelial cells cells from said suspension; and
 - (c) performing fluorescence activated cell sorting utilizing said antibodies so as to remove mature hepatocytes, mature bile duct cells, endothelial cells and mesenchymal cells from said suspension and

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multiparametric performing fluorescence activated cell sorting on said suspension utilizing antibody hepatic cell siđe marker, а forward scatter scatter, autofluorescence such that the cells said suspension in remaining isolated hepatoblasts.

- 14. The method of Claim 13 wherein the 10 antibody to a hepatic cell marker is monoclonal antibody 374.3.
 - 15. The method of Claim 13 wherein the hepatic cell marker is OC.3.
- 16. The method of Claim 13 wherein the single 15 cell suspension contains an agent capable of removing calcium from the surface of liver cells.
 - 17. The method of Claim 13 wherein the single cell suspension contains EGTA.
- 18. The method of Claim 13 wherein the single 20 cell suspension contains an enzyme capable of dissociating adult liver cells.
 - 19. The method of Claim 13 wherein the single cell suspension contains collagenase.
- 20. The method of Claim 13 wherein the single 25 cell suspension is chilled.
 - 21. The method of Claim 13 wherein the single cell suspension is at a temperature of between about 2 and 20°C.
- 22. Hepatocytes isolated by the method of 30 Claim 13.
 - 23. A method of treating liver dysfunction comprising the administration of hepatoblasts.
- 24. The method of Claim 23 wherein the administration comprises injecting said hepatoblasts into the liver via a vascular vessel.

- 25. The method of Claim 23 wherein the administration of comprises injecting said hepatoblasts into an ectopic site.
- 26. The method of Claim 23 wherein the administration comprises injecting said hepatoblasts into an ectopic site of the spleen.
 - 27. The method of Claim 23 wherein the hepatoblasts are isolated by the method of Claim 1.
- 28. The method of Claim 23 wherein the 10 hepatoblasts are isolated by the method of Claim 13.
 - 29. A method of forming an artificial liver comprising the utilization of hepatoblasts with a bioreactor.
- 30. The method of Claim 29 wherein the 15 hepatoblasts are isolated by the method of Claim 1.
 - 31. The method of Claim 29 wherein the hepatoblasts are isolated by the method of Claim 13.
 - 32. A method of forming an artificial liver comprising the utilization of hepatoblasts in a culture apparatus.
 - 33. The method of Claim 32 wherein the hepatoblasts are isolated by the method of Claim 1.
 - 34. The method of Claim 32 wherein the hepatoblasts are isolated by the method of Claim 13.

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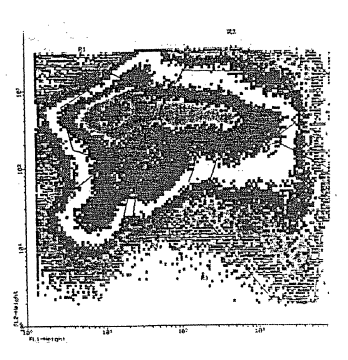


FIG.IA

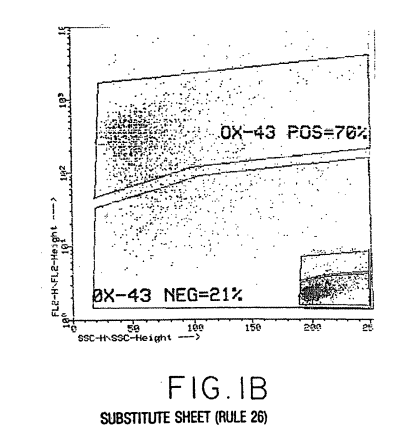


FIG. 1B SUBSTITUTE SHEET (RULE 26)

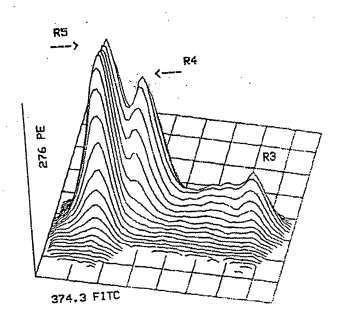


FIG. IC

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OX-43 OX-43

*

FIG.2A

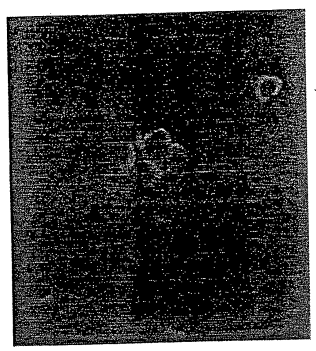


FIG.2B

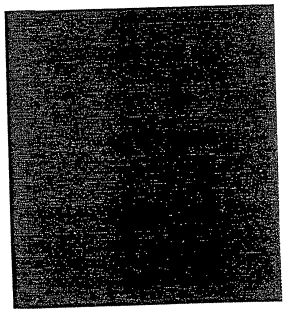


FIG. 2C SUBSTITUTE SHEET (RULE 26)

R3 R4 R5

Albumin



Serglycin

FIG.3

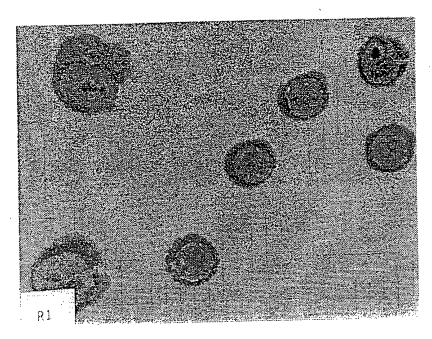


FIG. 4A

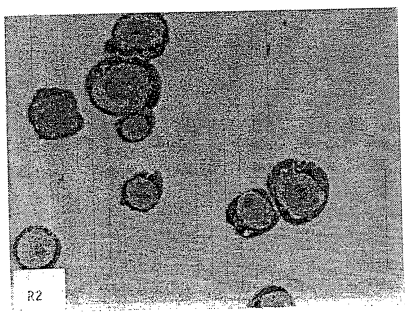


FIG. 4B

SUBSTITUTE SHEET (RULE 26)

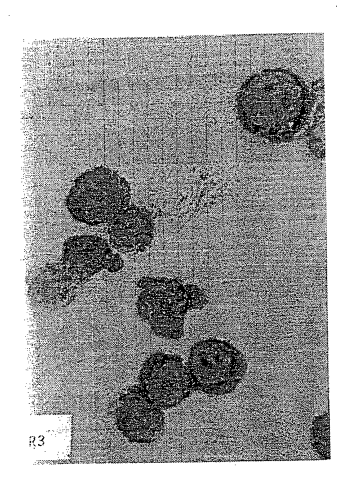


FIG. 4C

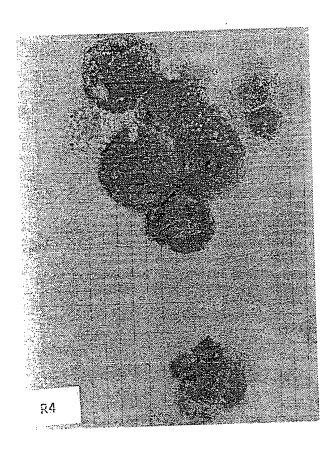


FIG. 4D

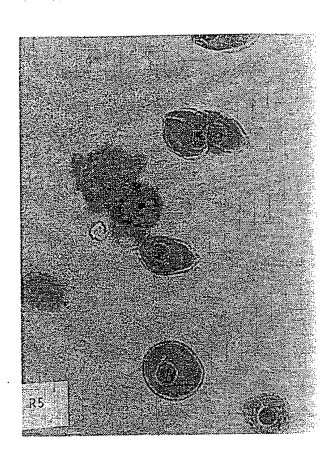


FIG.4E

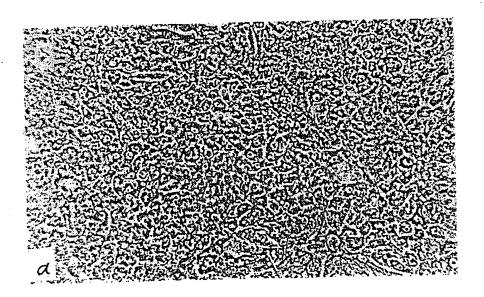


FIG. 5A

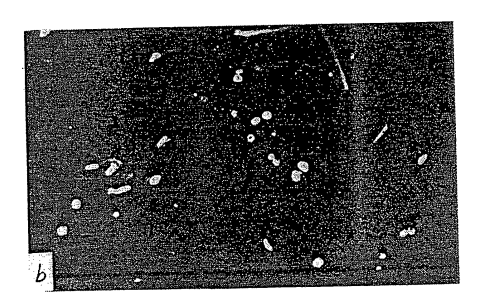


FIG. 5B

SUBSTITUTE SHEET (RULE 26)

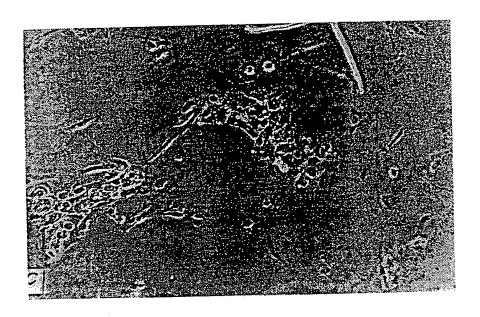


FIG. 5C

PCT/US94/13216 WO 95/13697

12/17 Flow Diagram of Hepatoblast Enrichment

Livers (8-9) mgs)

↓ Dispersion with EGTA and

then collagenase

Single Cell Suspension Preparation: Collagenase, EGTA, 4°C

₹ 10⁷ cells/8 mgs liver

♦ 3.2 ± 1.3% are ALB

↓ 2.5 ± 0.7% are AFP

+ 87.9 ± 2.5% are OX43/44*

Panning Red Blood Cell Panning (2X)

₹ 29 ± 5% of cells remain

+ 9.5 ± 1.2% are ALB

+ 9.8 ± 0.9% are AFP+

+ 80.4 ± 3.9% are OX43/44*

OX-43/OX-44 Panning (myeioid and endothelial cells)

₹ 16 ± 4% of cells remain

+ 14.8 ± 3.6% are ALB

14.9 ± 2.5% are AFP⁺

+ 69 ± 10% are OX43/OX44*

Fluorescence Activated Cell Sorting Negatively Sort for Contaminant Cell Populations:

OX-43 (CD)/OX-44 (CD37) Cells = precursors and mature forms of hemopoietic cells (myeloid, erythroid) and endothelial cells

Of remaining cells (OX-43 + OX-44 cells), sort for cells varying in OC.3 expression and granularity:

OX-43/(CD)/OX-44(CD37) Cells = mostly hepatic precursors, some residual hemopoiétic cell contaminants, stromal cells

OC.3⁺, granular cells = committed bile duct precursors (AFP⁺,ALB

OC.3, granular cells = committed hepatocyte precursors (AFP,ALB***)

OC.3⁺, agranular cells = early hepatoblasts (AFP⁺⁺⁺, albumin⁺and CK19⁻)

FIG. 6

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/3//7

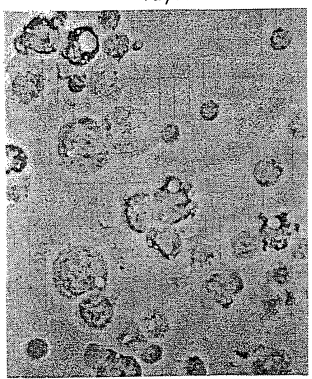


FIG.7A

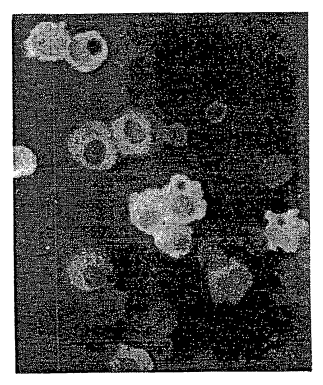


FIG. 7B SUBSTITUTE SHEET (RULE 26)

14/17

Original suspension
Panned cells

Original suspension
Panned cells

FIG.8

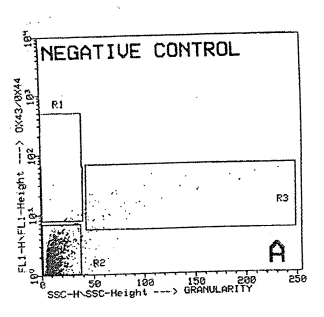


FIG. 9A

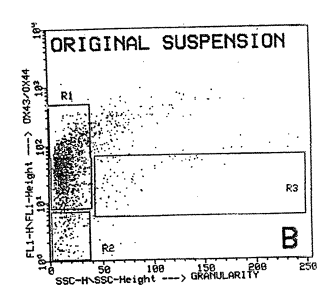


FIG. 9B SUBSTITUTE SHEET (RULE 26)

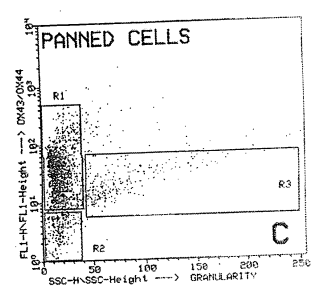


FIG.9C



FIG.10

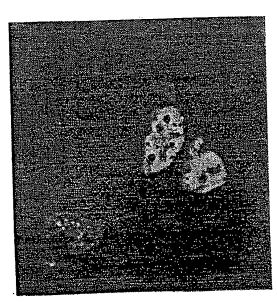


FIG. | | SUBSTITUTE SHEET (RULE 26)

International application No. PCT/US94/13216

A. CLASSIFICATION OF SUBJECT MATTER IPC(6) :A01N 1/02; G01N 33/533, 33/536, 33/538, 33/543, US CL :435/2, 7.21; 436/172, 175, 177, 178, 518, 536, 546 According to International Patent Classification (IPC) or to both	33/577 , 548 national classification and IPC			
DO CEANCITED				
B. FIELDS SEARCHED Minimum documentation searched (classification system followed) Minimum documentation searched (classification system followed)	. 548			
U.S. : 435/2, 7.21; 436/172, 175, 177, 178, 518, 536, 546,	aluded in the fields assembled			
Documentation searched other than minimum documentation to the				
Electronic data base consulted during the international search (na APS, MEDLINE, BIOSIS, EMBASE search terms: hepatoblast, liver stem cell, ox-43(44), treatment, therapy, bioreactor	oc.3, flow cytometry, liver dysfunction, artificial liver,			
C. DOCUMENTS CONSIDERED TO BE RELEVANT				
Category* Citation of document, with indication, where a				
l . wo of Homopoletic	BLOOD, Vol. 61, No. 3, issued March 1983, HOANG, T., et al, "Separation of Hemopoietic Cells From Adult Mouse Marrow by Use of Monoclonal Antibodies", pages 580-588, see entire document.			
PATHOBIOLOGY, Vol. 58, issued "An Antigenic Portrait of the Live pages 65-74, see entire document	er during Carcinogonious /			
X Further documents are listed in the continuation of Box	C. See patent family annex.			
Special categories of cited documents: 'A' document defining the general state of the art which is not considered to be of particular relevance 'E' earlier document published on or after the international filing date 'L' document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other appears reason (as specified) 'O' document referring to an oral disclosure, use, exhibition or other means	To later document published after the international time due to the state and not in conflict with the application but cited to understand the principle or theory underlying the invention "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art			
P document published prior to the international filing date but later that the priority date claimed	Date of mailing of the international search report			
Date of the actual completion of the international search				
21 FEBRUARY 1995	06 MAR 1995			
Name and mailing address of the ISA/US Commissioner of Patents and Trademarks Box PCT	NANCY J. PARSONS			
Washington, D.C. 20231	Telephone No. (703) 308-0196			

International application No. PCT/US94/13216

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	tion). DOCUMENTS CONSIDERED TO BE RELEVANT		n i va di No
	Citation of document, with indication, where appropriate, of the relevan	nt passages	Relevant to claim No.
Category*	CANCER RESEARCH, Vol. 48, issued 01 September 19 GERMAIN, L., et al, "Biliary Epithelial and Hepatocytheliage Relationships in Embryonic Rat Liver as Determinent the Differential Expression of Cytokeratins, alpha-Fetophalbumin, and Cell Surface-exposed Components", page 4918, see entire document.	1988, ic Cell mined by protein,	1-22, 27, 28, 30, 31, 33, 34
Y	AMERICAN JOURNAL OF PHYSIOLOGY, Vol. 263 1992, SIGAL, S.H., et al, "The liver as a stem cell and system", pages G139-G148, see entire document.		23-28
Y	IN VITRO CELL. DEV. BIOL., Vol. 29A, issued Malli, A.P., et al, "Culturing of Primary Hepatocytes as I Aggregates in a Packed Bed Bioreactor: A Potential Biological Primary, pages 249-254, see entire document.	rch 1993, Entrapped oartificial	29-34
	*		
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Form PCT/ISA/210 (continuation of second sheet)(July 1992)*

International application No. PCT/US94/13216

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
Box I Observations where certain claims were found unsearchand (continued of the following recents:
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2. Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This International Searching Authority found multiple inventions in this international application, as follows:
Please See Extra Sheet.
1. X As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment
of any additional fee. 3. As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:
4. No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark on Protest The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.

Form PCT/ISA/210 (continuation of first sheet(1))(July 1992)*

International application No. PCT/US94/13216

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING This ISA found multiple inventions as follows:

- 1. Claims 1-28, drawn to a method of isolating hepatoblasts, the hepatoblasts, and a method of treating liver dysfunction using hepatoblasts.
- II. Claims 29-34, drawn to a method of forming an artificial liver using hepatoblasts.

The claims of Groups I and II are not so linked by a special technical feature within the meaning of PCT Rule 13.2 so as to form a single inventive concept because the methods of Group I require different reagents and method steps and have different outcomes than the method of Group II. Additionally, the hepatoblasts of Group I have many different uses as shown by the two distinct methods of using them recited in Groups I and II.

Form PCT/ISA/210 (extra sheet)(July 1992)*

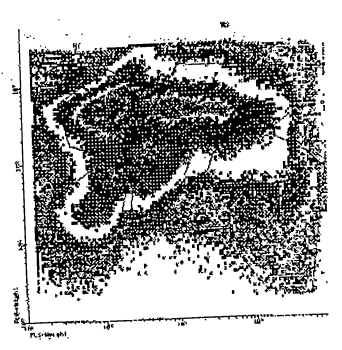


FIG.IA

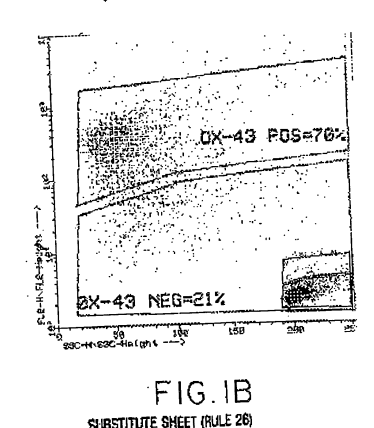


FIG. 1B SUBSTITUTE SHEET (RULE 26)

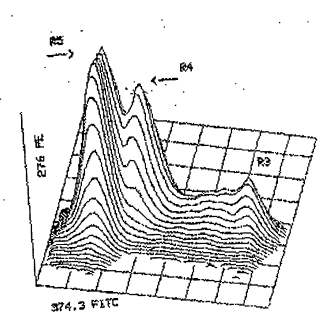


FIG. IC

3/17

OX-43

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A

FIG.2A

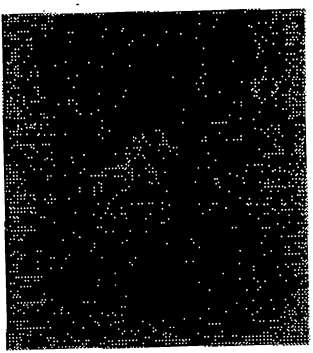


FIG.2B

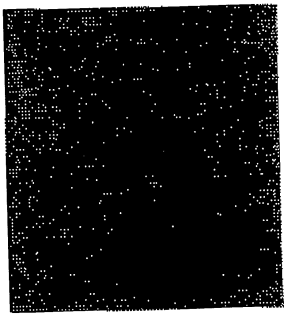


FIG.2C

SUBSTITUTE SHEET (RULE 26)

R3 R4 R5

Albumin



Serglycin

FIG.3

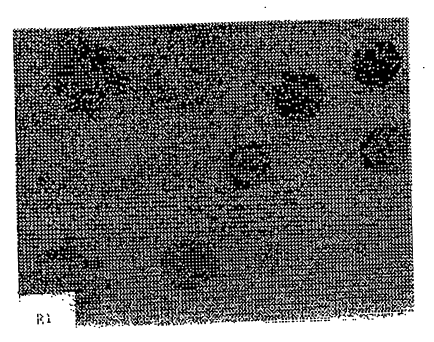


FIG. 4A

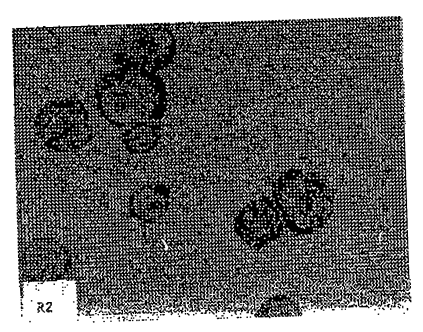


FIG. 4B

SUBSTITUTE SHEET (RULE 26)

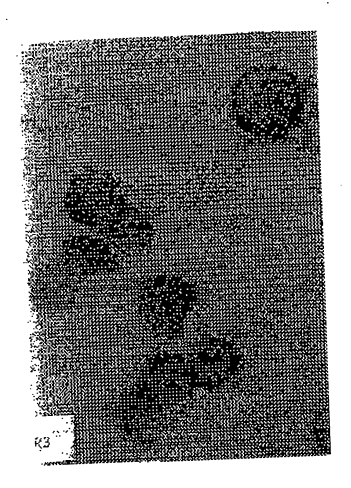


FIG. 4C

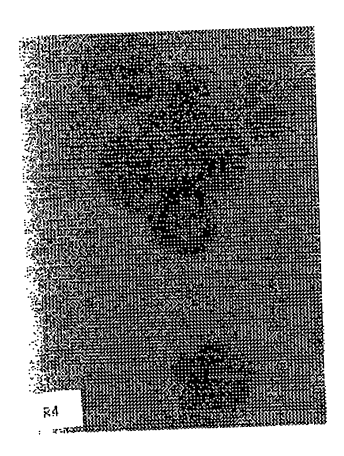


FIG. 4D

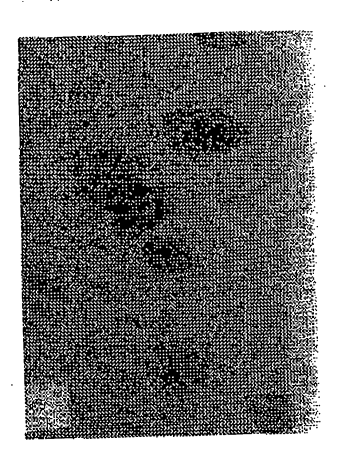


FIG.4E

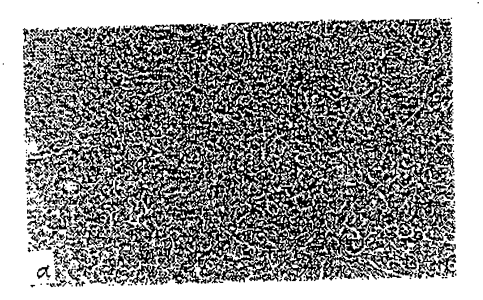


FIG. 5A

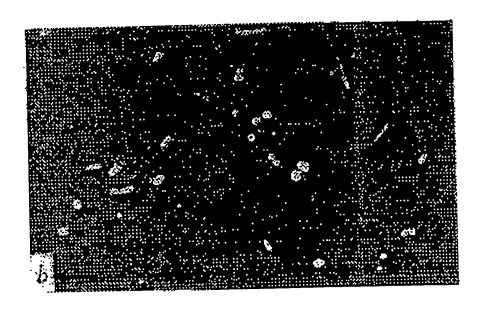


FIG. 5B

SUBSTITUTE SHEET (RULE 26)

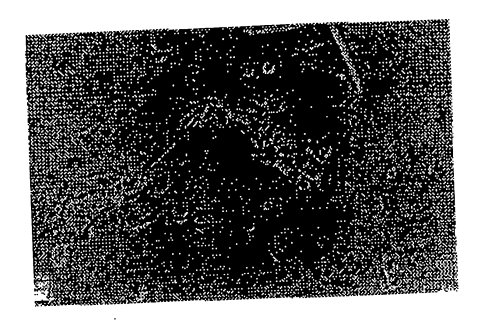


FIG. 5C

Flow Diagram of Hepatoblast Enrichment

Livers (8-9) mgs)

+ Dispersion with EGTA and

then collagenase

Single Cell Suspension Preparation: Collagenass. EGTA, 4°C

+ 107 cells/8 mgs liver

4 3.2 ± 1.3% are ALB

\$ 2.5 ± 0.7% are AFP'

+ 87.9 ± 2,5% are OX43/44*

Panning Red Blood Cell Panning (2X)

+ 29 ± 5% of cells remain

+ 9.5 ± 1.2% are ALB

\$ 9.8 ± 0.9% are AFP

+ 80.4 ± 3.9% are OX43/44*

OX-48/OX-44 Panning (myeloid and endothelial cells)

16 ± 4% of cells remain

+ 14.8 ± 3.6% are ALB+

₹ 14.9 ± 2.5% are AFP*

+ 69 ± 10% are OX43/OX44*

Fluorescence Activated Cell Sorting

Negatively Sort for Contaminant Cell Populations:

OX-43 (CD)/OX-44 (CD37) Cells = precursors and mature forms of hemopoietic cells (myeloid, erythroid) and endothelial cells

Of remaining cells (OX-43"+ OX-44" cells), sort for cells varying in OC.3 expression and granularity:

OX-48/(CD)/OX-44(CD37) Cells = mostly hepatic precursors, some residual hemopoietic cell contaminants, stromal cells

OC.3*, granular cells = committed bile duct precursors (AFP*,ALB

OC.3, granular cells = committed hepatocyte precursors (AFP',ALB'**)

OC.3*, agranular cells = early hepatoblasts (AFP***, albumin*and CK19⁻)

FIG. 6

/3//7

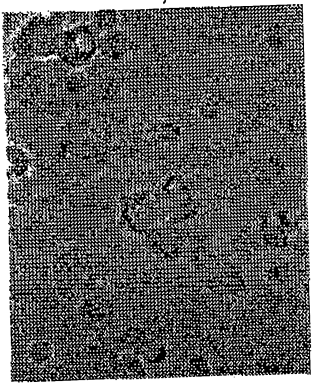


FIG. 7A



FIG. 7B SUBSTITUTE SHEET (RULE 26)

Original suspension
Panned cells

Original suspension
Panned cells

F1G.8

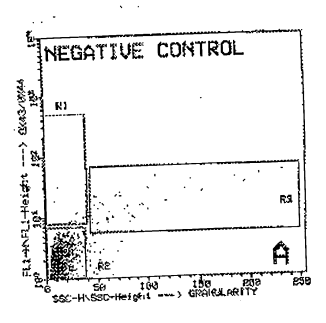


FIG. 9A

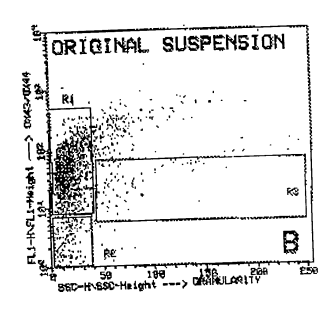


FIG. 9B SUBSTITUTE SHEET (RULE 26)

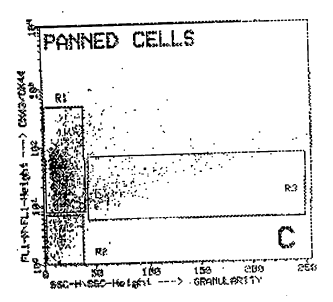


FIG.90

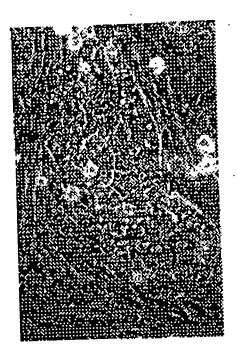


FIG. 10

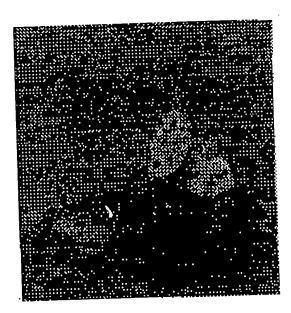


FIG. | | SUBSTITUTE SHEET (RULE 26)